less by the same arguments. In addition, the fraction of BO units changes in such a way as to enhance the growth rate as the PbO content increases, if BO is the structural unit necessary for growth. The scarcity of crystallizable species and the lower driving force for crystallization may account for the decreased growth rate observed in the 50.10 mol% PbO melt where the concentration of BO units would be less than for the other melts.

V. Conclusions

The determination of the temperature dependence of the growth rate of PbO-2B2O3 from several melts showed that the growth rate was greatly increased in those melts which contained slightly more than the stoichiometric quantity of PbO. This increase is attributed to the changes in the melt structure which occur in the interfacial region between the melt and the crystal. The concentration of boron atoms in fourfold coordination with oxygen atoms is increased as the PbO content is increased. These structural units are more readily incorporated into the crystal wherein all the boron atoms are in fourfold coordination with oxygen atoms. In addition, the fluidity of the melt increases with greater PbO concentration and hence the ease with which structural units may be oriented for attachment to the crystal is increased.

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I. Introduction and Previous Work

The crystallographic deformation mechanisms, fracture characteristics, and hardness relations for single crystals of MnTe and single-phase compositions in the system MnSe-MnTe were studied in the present work as a continuation of a study of the mechanical behavior of Mn (Group VI) solid solution compounds.

The stable forms of MnO, MnS, and MnSe have the fcc NaCl-type structure. In contrast, at normal temperatures MnTe has the hexagonal NiAs-type structure with a c/a ratio of 1.621. The NaCl and NiAs structures differ in the stacking of anion and cation layers. For the NaCl structure, the sequence is \( \text{AcBaAcB} \), whereas for the NiAs structure it is \( \text{AcBcAcBc} \). The capital letters represent anion layers and the lower case letters layers of the smaller cations. Manganese telluride appears to be very nearly stoichiometric, although Johansen' reported that MnTe transforms to the NaCl-type Structure above 1040°C and melts.

Deformation and Fracture of MnTe and MnSe-MnTe Solid Solutions

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Single crystals of MnTe and solid solutions in the system MnSe-MnTe were prepared; selected surfaces were indented with either a Vickers or a Knoop microindenter. Information on mechanical behavior was obtained by observation of slip traces and by study of Knoop hardness anisotropy. Hexagonal NiAs-type MnTe and MnTe-rich Mn(Te,Se) solid solutions show plastic deformation attributable to \( \{101\} \) twinning, basal slip, and pencil glide in \( \{1120\} \). Fracture occurs as primary \( \{0001\} \) and secondary \( \{1010\} \) cleavage. Vickers and Knoop hardnesses were greater on the basal plane than on prism planes. Cubic MnSe-rich Mn(Se,Te) solid solutions show both \( \{111\} \) and \( \{110\} \) \( \{110\} \) slip with \( \{100\} \) and \( \{110\} \) fracture in crushed fragments and around surface indentations. Knoop hardness anisotropy is like that in MnSe. The rate of solid solution hardening is greater than for comparable substitutions of sulfide ions in the MnSe matrix.

References


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at about 1170°C. It also undergoes an antiferromagnetic transformation on cooling below 310°K which results in a sharp contraction of the c axis.

Moore found that the primary deformation mechanism in MnO is slip in \( <110> \) directions on \( (110) \) planes, which is the usual mechanism in NaCl-type ionic crystals. Chao et al. also found that the \( (110) <110> \) mechanism was primary in MnS but detected secondary glide on \( (111) \) planes. Manganous selenide has \( (111) <110> \) and \( (111) <110> \) as the primary and secondary slip mechanisms, respectively. The change in primary mechanism in MnSe is related to a decrease in the ionic character of the chemical bond with increased anion size.

Fracture in MnO, MnS, and MnSe is characterized by easy \( (100) \) cleavage and secondary \( (110) \) fracture. Around Vickers microhardness indentations, MnS showed primarily \( (110) \) fracture because of multiple slip interactions on adjacent \( (110) \) slip planes, whereas interactions on \( (111) \) slip planes in MnSe produced \( (100) \) fracture.

### II. Experimental Procedure

(1) Preparation and Purification of Raw Materials

Manganous telluride and selenide were prepared from Mn powder made from tripoly vacuum-distilled electrolytic Mn and from Te or Se powder prepared from high-purity shot. The powders were mixed in appropriate proportions, placed in a vacuum-degassed graphite crucible, encapsulated in a borosilicate glass tube under vacuum, and reacted at 600°C for several days. Both MnTe and MnSe were purified further by zone refining.

Microscopy revealed no other phases except an occasional small inclusion which was apparently MnO. X-ray data gave average values of \( a_0=4.137 \) and \( c_0=6.706 \) A; average values of \( a_0=4.137 \) and \( c_0=6.706 \) A were reported in the literature by ten independent workers. The lattice parameter for MnSe was \( a_0=5.463 \) A; \( 5.462 \) A is the accepted value.

Chemical analyses of MnTe gave 30.2±0.3% Mn compared with 30.1% Mn for stoichiometry. For MnSe, analyses gave 40.8±0.4% Mn compared with 41.0% for stoichiometry.

(2) Crystal Preparation

Single crystals of MnTe, MnSe, and appropriate MnSe-MnTe compositions were prepared by a Bridgeman technique. The growth of high-quality single crystals of MnTe was difficult because of the polymorphic phase change at 1040°C and the c axis contraction at 310°K caused by the antiferromagnetic transformation. Usually a coarse-grained structure was obtained in which individual grains tended to be misoriented by up to 8° to 10°. This structure was a coarse mosaic in which all the mosaic parts were within ±5° of an average orientation. Often a “crystal” would show only two or three orientations differing by not more than ±3° from a mean value; these crystals were used whenever possible.

Single crystals were annealed for 12 h at 1000°C, oriented by the Laue back-reflection technique, and cut to reveal selected faces. For cubic crystals, (001), (011), and (111) planes were studied, whereas for hexagonal crystals (0001), (1010), and (1120) planes were selected. Cubic orientations were accurate to within 2° and hexagonal orientations to within about 8° to 10°.

Samples were mounted in Bakelite and mechanically polished using standard metallographic techniques. Care was taken during the latter stages of polishing to remove chemically any disturbed surface layers with an etchant of 1% HSO\(_4\), 1% H\(_3\)PO\(_4\), and 1% saturated oxalic acid solution. Subsequent Laue photographs showed no significant residual plastic deformation in the crystals.

Fig. 1. Vickers hardness impression on (0001) surface of MnTe crystal showing traces due to \( (1012) \) twinning (\( \times 250 \)).

### III. Results and Discussion

(1) Manganous Telluride

(A) Plastic Deformation: The modes of plastic deformation identified in MnTe were twinning on \( (1012) \) planes, slip involving the \( (0001) \) plane, and pencil glide with \( <1120> \) as the zone axis. The preferred mode varies with the exposed crystal plane and the stress direction.

Indentation into the basal plane produced patterns (Fig. 1) which may be attributed to \( (1012) \) twinning. The twin traces appear in two of the three possible \( <1120> \) intersection directions involving \( (1012) \) planes. (The activation of the twinning on only one side of the indenter reflects the previously explained slight misorientation of the \( (0001) \) plane.) Interference microscopy showed that this material was raised. This result further supports the twinning mechanism since, with \( c/a<\sqrt{3} \), \( (1012) \) twinning produces an elongation in the \( [0001] \) direction. Partridge and Roberts reported similar results in Mg single crystals which they identified as \( (1012) <110> \) twinning.

When either the \( (1120) \) or \( (1010) \) surface was indented with the indenter perpendicular to \( (0001) \), slip traces appeared perpendicular to \( (0001) \) (Figs. 2 and 3(A)). Movements on the basal, or \( (0001) \), plane must be present to account for both of these observations. However, each set of slip traces...
must also include movement with a [0001] vector to produce the total deformation. The planes which contain the latter displacement were not identified.

Figure 3(B) reveals wavy slip lines which must be attributed to many slip systems deforming simultaneously. This behavior suggests pencil glide, and since the <1120> Burgers vector has the shortest displacement distance, that possibility may be examined on a stereographic projection (Fig. 4). As shown on the (1010) projection, there are many possible planes within the common [1120] zone axis (and within the [2110] zone axis). These intersect the (1010) plane at a variety of angles, as is observed in Fig. 3(B). The same wavy deformation pattern is faintly visible on the (1120) surface to the left of the indenter in Fig. 2. Again pencil glide may account for this behavior from the [1210] and [2110] zones; however, with a steeper angle, the amount of slip should be less. This type of deformation may also be described as extensive cross-slip and is common for many materials, e.g. AgCl, AgBr, α-Fe, and Hg, all of which, like MnTe, are soft with easy dislocation movements.

(B) Fracture: The primary cleavage plane for MnTe appears in Fig. 2 and is (0001); this plane reveals bright shiny surfaces. A (1010) cleavage was also observed and it exposed a rather dull surface. Furthermore, pyramidal cleavage was found in some specimens around inclusions (Fig. 5). Evidently a moving (0001) cleavage crack meeting a foreign particle finds it energetically easier to move over and around some of the inclusions rather than through them.

(2) MnSe-MnTe System (MnTe-Rich Portion)

(A) Plastic Deformation and Fracture: Within the MnTe-rich single-phase region of the MnSe-MnTe system, samples of 9 and 19 at.% MnSe were studied. Both showed the same modes of plastic deformation found in pure MnTe: (1012) twinning, basal slip, and pencil glide in <1120>. Fracture
mechanisms were also the same as those in pure MnTe: (0001) and {1010} cleavage.

(B) Diamond Pyramid Hardness: Figure 6 shows the DPH results for the MnTe-rich solutions. The DPH on the basal plane was about 60% greater than that on (1010) for all compositions. The higher hardness of the basal plane reflects the difficulty of activating slip systems capable of moving material to the surface to make room for the indenter; this cannot be done by glide in $<1120>$ directions alone because they are all parallel to the (0001) plane.

(C) Knoop Hardness: Figure 7 shows the Knoop hardness results for these solutions. As with other single crystal specimens, Knoop hardness on the {1010} planes showed an anisotropy which depended on the orientation of the long axis of the indenter and the symmetry of the slip systems of the crystal. All orientations of the indenter on the basal plane, however, gave the same hardness within the ±4% limits of experimental error, but, as with Vickers hardness, the basal plane was much harder than the (1010) plane. The variation on the (1010) plane reflects the many slip systems which can be activated during pencil glide for the [0001] indenter orientation but which cannot be supplied by the predominantly basal slip found at the [1210] indenter orientation (Fig. 3).

Similar tests on the (1120) plane gave results virtually identical to those found on (1010).

(3) MnSe-MnTe System (MnSe-Rich Portion)

(A) Plastic Deformation and Fracture: Sodium chloride-type MnSe has $<111><110>$ and $<110><110>$ as primary and secondary slip mechanisms, respectively. Within the cubic single-phase region, the compositions studied were 93 at.% MnSe-7 at.% MnTe and 87 at.% MnSe-13 at.% MnTe. Both compositions showed slip traces due to $<111><110>$ and $<110><110>$ glide, with the amount of $<110>$ glide somewhat more extensive than for comparable substitutions of sulfide for selenide ions. Figure 8 shows slip traces due to both slip mechanisms on the (001) plane of a 93 MnSe-7MnTe crystal. The (110) slip traces form a square arrangement in $<100>$ directions, whereas the fainter (111) traces are at 45° to $<100>$ directions. Similar indentations on the (011) and (111) planes confirmed these findings.
indentations and decreased the relative incidence of (100)\textsubscript{\(P\)} fractures. The (110)\textsubscript{\(P\)} fracture is a consequence of dislocation interactions on intersecting (110)\textsubscript{\(P\)} slip planes,\(^{18,19}\) whereas the (100)\textsubscript{\(P\)} fracture stems from dislocation interactions on adjacent (111) slip planes.\(^{20}\)

(B) Diamond Pyramid Hardness: Although the Vickers hardness was essentially the same on the (001), (011), and (111) planes, the values were higher than those observed for comparable substitutions of sulfide for selenide ions (dashed lines in Fig. 6). The difference in ionic radii between S\textsuperscript{2-} and Se\textsuperscript{2-} (1.74 vs 1.91 Å, Goldschmidt) is 8.9\%, whereas that between Te\textsuperscript{2-} and Se\textsuperscript{2-} (2.11 vs 1.91 Å) is about 10.5\%. Although the bonding is not totally ionic, and ionic radii, therefore, are not strictly applicable, this comparison indicates that a slightly larger size mismatch in the case of the telluride may cause the observed higher rate of hardening.

(C) Knoop Hardness: The Knoop hardness results for both compositions followed the same pattern as for pure MnSe; that is, for indentations on the (001) plane, the hardness was lowest when the long axis of the indenter was parallel to \(<110>\) (Fig. 7). Also, like the Vickers hardness results, the rate of hardening was greater than that found for the substitution of sulfide for selenide ions. Similar results were found on the (011) plane.

IV. Conclusions

(1) Manganous telluride shows patterns consistent with three modes of plastic deformation around Vickers and Knoop indentations: \(\{10\overline{1}2\}\) twinning, basal slip, and pencil glide in \(<11\overline{2}0>\).

(2) Fracture in MnTe occurs as primary (0001) and secondary (1010) cleavage both in crushed fragments and around surface indentations. Pyramidal cleavage can also occur, especially around small inclusions.

(3) Solid solutions formed by substitution of selenide ions in the MnTe matrix show the same plastic deformation and fracture characteristics as MnTe. The basal plane is harder than the prism planes, reflecting the difficulty of activating enough independent slip systems by basal indentation.

(4) The addition of telluride ions to MnSe results in an early appearance of (110) slip and produces a slightly higher rate of solid-solution hardening than a comparable substitution of sulfide ions.

References


Discussion and Notes

Comments on "Wetting of Binary Aluminum Alloys in Contact with Be, B₄C, and Graphite"

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Recently Manning and Gurganus' reported measurements of the wettability of solid Be, B₄C, and graphite with various liquid alloys of Al. Some comments on their interpretation of these measurements are offered.

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